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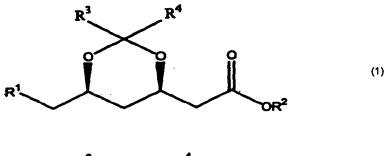
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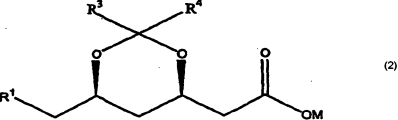
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(54) Title: PROCESS FOR THE PREPARATION OF DIOXANE ACETIC ACID ESTERS





(57) Abstract: Process for the preparation of an ester of formula (1), wherein R¹represents a leaving group, CN, OH or a COOR⁵ group, R³ and R⁴each independently represent a 1-3 C alkyl group, and R² and R⁵ each independently represent an ester residue, wherein the corresponding salt with formula (2), wherein M represents H or an alkali (earth) metal in an inert solvent is contacted with an acid chloride forming agent to form the corresponding acid chloride, and the acid chloride is contacted with an alcohol with formula R²OH in the presence of N-methyl-morpholine. Preferably M represents an alkali metal, and R² represents an alkyl group, particularly a t-butyl group. (1), (2)

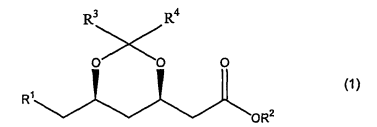


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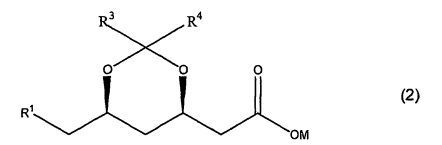
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PROCESS FOR THE PREPARATION OF DIOXANE ACETIC ACID ESTERS

5 The invention relates to a process for the preparation of an ester of formula (1)



wherein R¹ represents a leaving group, CN, OH or a COOR⁵ group, R³ and R⁴ each independently represent a C1-3 alkyl group and R² and R⁵ each independently represent an ester residue, wherein the corresponding salt with formula (2)



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wherein M represents H or an alkali (earth) metal in an inert solvent is contacted with an acid chloride forming agent to form the corresponding acid chloride, and the acid chloride is contacted with an alcohol with formula R²OH in the presence of N-methyl morpholine (NMM).

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Many processes for the preparation of esters are known in the art, for instance the preparation of esters via the formation of the acid chloride. It was, however, to be expected that such processes would not lead to high yields due to the lack of stability of the present compound under acidic conditions.

It is the object of the invention to provide a process for the

preparation of esters with high yield in a robust process, even at large scale and with relatively high concentrations.

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Surprisingly it has been found that even sterically hindered esters that are difficult to obtain in esterifications like t-butyl esters of the acid unstable molecules of formula (1), can be obtained in high yield in an easily reproducible process.

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With the process according to the invention esters with formula (1) that are unstable under acidic conditions, for instance with pH < 4, can be prepared in high yield.

R¹ represents a leaving group, CN, OH or a COOR⁵ group wherein R⁵ represents an ester residue, for example an alkyl group with for instance 1-6 C-atoms, or an aryl group with for instance 6-12 C-atoms. A leaving group by definition is a group that can easily be replaced, for example a halogen, for instance Cl, Br or I; a tosylate group; a mesylate group; an acyloxy group, with, for instance, 1-6 C-atoms in particular an acetoxy group; a phenacetyloxy group; an alkoxy group with, for instance, 1-6 C-atoms or an (hetero) aryloxy group with, for instance, 6-12 C-atoms. Preferably R¹ represents Cl.

R² represents an ester residue, preferably an alkyl group, for instance an alkyl group with 1-6 C-atoms or an aryl group, for instance an aryl group with 6-12 C-atoms, in particular a methyl, ethyl, propyl, isobutyl or t.butyl group. An important group of esters that can be prepared with the process according to the invention are t.butyl esters.

R³ and R⁴ each independently represent a C1-C3 alkyl group, for instance a methyl or ethyl group. Preferably R³ and R⁴ both represent methyl.

M in formula (2) can be chosen from the group of H, alkali metals, for instance lithium, sodium, potassium and alkali earth metals, for instance magnesium or calcium. Preferably M represents sodium or potassium.

The acid chloride forming agent can be chosen from the group of reagents that is generally known as such. Suitable examples of acid chloride forming agents are oxally chloride, thionyl chloride, PCl₃, PCl₅, and POCl₃. Preferably the acid chloride forming agent is used in an excess relative to the amount the salt with formula (2), for instance between 1 and 3 equivalents, more preferably between 1.2 and 1.8 equivalents.

If desired, in the acid chloride formation also a catalyst may be present. The amount of catalyst may for instance vary from 0-1, preferably 0-0.5 equivalents, calculated with respect to the amount of salt with formula (2). Higher

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amounts of catalyst are also possible, but will normally have no extra advantageous effect. Preferably the amount of catalyst, if any, will be between 0.05 and 0.2 equivalents calculated with respect to the salt with formula (2). Suitable catalysts are the catalysts generally know to accelerate acid chloride formation, for instance dimethylformamide (DMF) and N-methylpyrrolidone (NMP).

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The conversion of the acid chloride into the ester with formula (1) is carried out in the presence of an alcohol with formula R²OH. The amount of alcohol with formula R²OH is not very critical and preferably is between 1 and 15 equivalent calculated with respect to the amount of salt with formula (2), more preferably between 2 and 13, most preferably between 3 and 6. Surprisingly it has been found that even t.-butyl esters can be prepared with high yield using a relatively low amount of t.-butyl alcohol.

The conversion of the acid chloride into the ester with formula (1) is carried out in the presence of NMM. In practice a small amount of NMM, efficient to catch eventually remaining free HCl, for instance 1.5 to 2.5, preferably 1.8 to 2.0 equivalents calculated with respect to the amount of salt with formula (2) is applied. When a large excess of acid chloride forming agent is used, preferably higher amounts of NMM are used, and when a lower excess of acid chloride forming agent is used, preferably lower amounts of NMM are used.

The acid chloride formation reaction preferably is carried out at a temperature between -30° and 60°C, more preferably between 20 and 50°C. The conversion of the acid chloride into the ester with formula (1) preferably is carried out at a temperature between 20 and 80°C, more preferably between 20 and 50°C.

The process of the present invention may be carried out in one step. Preferably first the salt with formula (2) is converted into the corresponding acid chloride, and subsequently the acid chloride is contacted with the alcohol with formula R²OH and NMM. In a particularly preferred embodiment the acid chloride formed is quenched with NMM and the alcohol with formula R²OH.

The product with formula 1, wherein R¹ represents a leaving group may subsequently be converted into the corresponding compound wherein R¹ represents an acyloxy group. This can be achieved in a manner known per se, for instance by reaction with an acyloxylating agent for instance a carboxylic or sulphonic acid, a quaternary ammonium or phosphonium salt, a carboxylic or sulphonic acid quaternary ammonium or phosphonium salt or a combination thereof. Preferably a combination of a quaternary phosphonium salt and a carboxylic or sulphonic acid salt is used as the acyloxylating agent.

Subsequently the compound with formula 1, wherein R¹ represents an acyloxy group can be converted in the corresponding compound wherein R¹ represents a hydroxy group, for instance by subjecting it to solvolysis in the presence of a base. Suitable bases are, for instance, alkali (earth) metal hydroxides or carbonates or organic bases, for instance alkali (earth) metal carboxylic acids, for instance acetates, ammonia, pyridines, amines, for instance triethylamine and the like. The invention will be elucidated by the following examples.

Example I

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10 1864 g of an aqueous solution of the (4R-cis)-(6-chloromethyl)-2,2-dimethyl-1,3-dioxane-4-yl-acetic acid, sodium salt (3.31 moles) and 4.8 L of toluene were mixed and water was removed by azeotropic distillation under reduced pressure. Subsequently, 870 g of fresh toluene were added and removed by distillation. To the obtained suspension was added 33.4g of NMP. Then 588 g of oxalyl chloride were added while maintaining the temperature at 20 °C. The resulting mixture was stirred for 6 hours at 20-25 °C and then slowly added to a mixture of 979 g of t.-butanol and 836 g of *N*-methyl morpholine. After stirring for 1 hour, 3966 g of an 8% (w/w) aqueous NaOH solution was added and the resulting mixture stirred for 1.5 hours at 40 °C. After washing the organic phase with 3300 g of water, 3064 g of a toluene solution of the desired t.-butyl ester was obtained, corresponding to 751 g (81%) of product.

Example II

In a 100 ml HEL Vessel with 4 blade stirrer 8.0 g (4R-cis)-(6-chloromethyl)-2,2-dimethyl-1,3-dioxane-4-yl-acetic acid, sodium salt (92.4%; 30 mmol) was suspended in 41 g toluene and 0.3 g NMP (3 mmol). In 1h 4.5 g (36 mmol) oxalylchloride was dosed at a temperature of 15-20°C. The reaction mixture (50 g) was stirred for 2.5 hours. The reaction mixture was split into 2 parts: Part A (23.83 g) and part B (24.25 g). Part A of the reaction mixture was dosed during 1 h. to a mixture of 22.2 g (20 eq.) t.-butanol and 3.0 g (2 eq.) NMM at 25°C. The reaction mixture was stirred overnight and analyzed by GC. The yield of the t.-butyl ester was 88%.

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Examples III-V

Following the same procedure as described in Example I, the ethyl, isopropyl and n-hexyl esters, respectively, are prepared wherein instead of 4 eq. butanol, now 4 eq. ethanol, 4 eq. isopropanol and 4 eq. n-hexanol, respectively is used. The yield of the desired ethyl, isopropyl and n-hexyl ester was 89 mol%, 88 mol% and 84 mol% respectively, calculated with respect to the sodium salt starting material.

Example VI

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A mixture of 35.0 g of t-butyl (4R-cis)-(6-chloromethyl)-2,2-dimethyl-1,3-dioxane-4-yl-acetate, 14.8 g of tetrabutyl phosphonium bromide, 16.0 g of potassium acetate and 5.9 g of toluene were heated to 105 °C under reduced pressure. After 22 hours at this temperature the reaction mixture was cooled to ambient temperature after which 400 g of heptane and 350 g of water were added. The organic phase was washed with 150 g of water and subsequently treated with 3.0 g of activated carbon. After filtration of the carbon, the solution was concentrated and subsequently cooled to -10 °C after which crystallised product was isolated by means of filtration. Yield 24.9 g (76%) of a white crystalline material

CLAIMS

1. Process for the preparation of an ester of formula (1),

$$\mathbb{R}^{3}$$

$$\mathbb{R}^{4}$$

$$\mathbb{R}^{1}$$

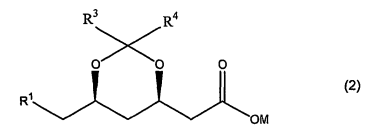
$$\mathbb{R}^{2}$$

$$\mathbb{R}^{2}$$

$$\mathbb{R}^{3}$$

$$\mathbb{R}^{4}$$

wherein R¹ represents a leaving group, CN, OH or a COOR⁵ group, R³ and R⁴ each independently represent a 1-3 C alkyl group, and R² and R⁵ each independently represent an ester residue, wherein the corresponding salt with formula (2),



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wherein M represents H or an alkali (earth) metal in an inert solvent is contacted with an acid chloride forming agent to form the corresponding acid chloride, and the acid chloride is contacted with an alcohol with formula R²OH in the presence of N-methyl-morpholine.

- 20 2. Process according to claim 1, wherein M represents an alkali metal.
 - 3. Process according to claim 1 or 2, wherein R² represents an alkyl group.
 - 4. Process according to claim 3, wherein R² represents a t.-butyl group.
 - 5. Process according to any one of claims 1-4, wherein the acid chloride forming agent is oxalylchloride.
- 25 6. Process according to any one of claims 1-5, wherein the acid chloride formation is performed in the presence of a catalyst.
 - 7. Process according to any one of claims 1-6, wherein the amount of alcohol

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with formula R²OH is between 3 and 6 equivalents calculated with respect to the amount of salt with formula (2).

8. Process according to any one of claims 1-7, wherein first the salt with formula (2) is converted into the corresponding acid chloride and subsequently the acid chloride is contacted with the alcohol with formula R²OH and N-methyl-morpholine.

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- 9. Process according to claim 8, wherein the acid chloride is quenched with the alcohol with formula R²OH and N-methyl-morpholine.
- 10. Process according to any one of claims 1-9, wherein R¹ represents a leaving group, and wherein the ester of formula 1 wherein R¹ represents a leaving group is subsequently converted into the corresponding ester with formula 1 wherein R1 represents an acyloxy group.
- 11. Process according to claim 10, wherein first an ester of formula 1 wherein R¹ represents an acyloxy group is prepared and subsequently the ester of formula 1 is converted into the corresponding compound with formula 1 wherein R¹ represents OH.

	INTERNATIONAL SEARCH REPO	RT		Application No	
			PCT/NL	03/00435	
A. CLASSI IPC 7	FIGATION OF SUBJECT MATTER CO7D319/06				
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	SEARCHED				
IPC 7	ocumentation searched (classification system followed by classificat ${\tt C070}$	ton symbols)			
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	lata base consulted during the International search (name of data be ternal, WPI Data, PAJ, CHEM ABS Data	•	1, search terms u	sed)	
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT				
Category *	Citation of document, with indication, where appropriate, of the re	levant passages		'Relevant to claim No.	
х	WO 02 06266 A (MINK DANIEL ;DSM KOOISTRA JACOB HERMANUS MATTHE (24 January 2002 (2002-01-24) page 5, line 4 -page 6, line 31;	1-11			
	example VII				
A	WO 00 68221 A (EGYT GYOGYSZERVEG GYAR ;VERECZKEYNE DONATH GYOERGY: 16 November 2000 (2000-11-16) page 7, paragraph 4	1-11			
A	GB 885 516 A (ARTHUR HENRY CLARKS 28 December 1961 (1961-12-28) page 1: left-hand column, paragra page 2: right-hand column, last p	1-11			
		-/- -			
X Furti	her documents are listed in the continuation of box C.	Patent family	members are lis	ted in annex.	
Special categories of cited documents: A document defining the general state of the art which is not considered to be of particular retevance E earther document but published on or after the international filling date L document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) O document referring to an oral disclosure, use, exhibition or other means P document published prior to the international filling date but later than the priority date claimed		"Y" later document published after the International filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family			
	actual completion of the international search 6 August 2003	Date of mailing of 12/09/2		search report	
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	European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fex: (+31-70) 340-3016	Stroeter, T			

INTERNATIONAL SEARCH REPORT

Internat Application No
PCT/NL 03/00435

Citedony* Citation of document, with indication, where appropriate, of the relevant passages A MARCH J: "ALIPHATIC NUCLEOPHILIC SUBSTITUTION" ADVANCED ORGANIC CHEMISTRY. REACTIONS, MECHANISMS, AND STRUCTURE, NEW YORK, JOHN WILEY & SONS, US, 1992, page 392 XP002217003 ISBN: 0-471-60180-2 chapter 0-20 page 392 A MURPHY C F, KOEHLER R. E.: "CHEMISTRY OF CEPHALOSPORIN ANTIBIOTICS. XVIII." J. ORG. CHEM., vol. 35, no. 7, 1970, pages 2429-2430, XP002252465 right—hand column, first paragraph Hetevant to cialm No. 1-11		INION DOCUMENTS CONSIDERED TO BE BEI EVANT	
A MARCH J: "ALIPHATIC NUCLEOPHILIC SUBSTITUTION" ADVANCED ORGANIC CHEMISTRY. REACTIONS, MECHANISMS, AND STRUCTURE, NEW YORK, JOHN WILEY & SONS, US, 1992, page 392 XP002217003 ISBN: 0-471-60180-2 chapter 0-20 page 392 A MURPHY C F, KOEHLER R. E.: "CHEMISTRY OF CEPHALOSPORIN ANTIBIOTICS. XVIII." J. ORG. CHEM., vol. 35, no. 7, 1970, pages 2429-2430, XP002252465	Jategory °		
SUBSTITUTION" ADVANCED ORGANIC CHEMISTRY. REACTIONS, MECHANISMS, AND STRUCTURE, NEW YORK, JOHN WILEY & SONS, US, 1992, page 392 XP002217003 ISBN: 0-471-60180-2 chapter 0-20 page 392 A MURPHY C F, KOEHLER R. E.: "CHEMISTRY OF CEPHALOSPORIN ANTIBIOTICS. XVIII." J. ORG. CHEM., vol. 35, no. 7, 1970, pages 2429-2430, XP002252465		Cuation of document, with indication, where appropriate, of the relevant passages	Retevant to claim No.
CEPHALOSPORIN ANTIBIOTICS. XVIII." J. ORG. CHEM., vol. 35, no. 7, 1970, pages 2429-2430, XP002252465	4	SUBSTITUTION" ADVANCED ORGANIC CHEMISTRY. REACTIONS, MECHANISMS, AND STRUCTURE, NEW YORK, JOHN WILEY & SONS, US, 1992, page 392 XP002217003 ISBN: 0-471-60180-2 chapter 0-20	1-11
	A	CEPHALOSPORIN ANTIBIOTICS. XVIII." J. ORG. CHEM., vol. 35, no. 7, 1970, pages 2429-2430, XP002252465	1-11

INTERNATIONAL SEARCH REPORT

Internal application No PCT/NL 03/00435

Patent document cited in search report		Publication date		Patent family member(s)	Publication date
WO 0206266	A	24-01-2002	NL	1015744 C2	22-01-2002
			ΑU	7583001 A	30-01-2002
			BR	0112535 A	01-07-2003
			CA	2415963 Al	24-01-2002
			CZ	20030163 A3	14-05-2003
			EP	1317440 A1	11-06-2003
			WO	0206266 A1	24-01-2002
			NO	20030025 A	03-01-2003
WO 0068221	Α	16-11-2000	HU	9901526 A2	28-04-2001
			AU	4600200 A	21-11-2000
			CA	2373077 A1	16-11-2000
			CN	1349522 T	15-05-2002
			CZ	20013965 A3	17-04-2002
			EP	1178980 A1	13-02-2002
			HR	20010846 A1	28-02-2003
			WO	0068221 A1	16-11-2000
			JР	2002544207 T	24-12-2002
			PL	351145 A1	24-03-2003
			SK	15842001 A3	04-04-2002
GB 885516	Α	28-12-1961	NONE	، میروری چانند بوبدوری کا سام اساس است.	